C:\stnweb\Queries\10618288-2.str Part II - Broader Guerg 12 chain nodes : 7 8 9 10 11 12 13 14 16

```
7 8 9 10 11 12 13 14 16

ring nodes:
    1 2 3 4 5 6

chain bonds:
    2-7 5-16 7-8 7-9 7-10 10-11 11-12 11-13 13-14

ring bonds:
    1-2 1-6 2-3 3-4 4-5 5-6

exact/norm bonds:
    1-2 1-6 2-3 2-7 3-4 4-5 5-6 7-8 7-9 7-10 10-11 11-12 11-13
    13-14

exact bonds:
    5-16

isolated ring systems:
    containing 1:
```

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:CLASS 8:CLASS 9:CLASS

10:Atom 11:CLASS 12:CLASS 13:CLASS 14:CLASS 16:Atom

```
Welcome to STN International
NEWS
                Web Page URLs for STN Seminar Schedule - N. America
                 "Ask CAS" for self-help around the clock
NEWS
                PATDPAFULL - New display fields provide for legal status
NEWS 3
        FEB 28
                data from INPADOC
NEWS 4 FEB 28 BABS - Current-awareness alerts (SDIs) available
NEWS 5 MAR 02 GBFULL: New full-text patent database on STN
NEWS 6 MAR 03 REGISTRY/ZREGISTRY - Sequence annotations enhanced
NEWS 7 MAR 03 MEDLINE file segment of TOXCENTER reloaded
NEWS 8 MAR 22 KOREAPAT now updated monthly; patent information enhanced
    9 MAR 22 Original IDE display format returns to REGISTRY/ZREGISTRY
NEWS
NEWS 10 MAR 22 PATDPASPC - New patent database available
NEWS
     11 MAR 22 REGISTRY/ZREGISTRY enhanced with experimental property tags
NEWS 12 APR 04 EPFULL enhanced with additional patent information and new
                fields
NEWS 13 APR 04 EMBASE - Database reloaded and enhanced
NEWS 14 APR 18 New CAS Information Use Policies available online
NEWS 15 APR 25 Patent searching, including current-awareness alerts (SDIs),
                based on application date in CA/CAplus and USPATFULL/USPAT2
                may be affected by a change in filing date for U.S.
                applications.
NEWS
     16 APR 28
                Improved searching of U.S. Patent Classifications for
                U.S. patent records in CA/CAplus
     17 MAY 23
                GBFULL enhanced with patent drawing images
NEWS
     18 MAY 23
                REGISTRY has been enhanced with source information from
                CHEMCATS
NEWS 19 JUN 06 STN Patent Forums to be held in June 2005
NEWS 20 JUN 06 The Analysis Edition of STN Express with Discover!
                 (Version 8.0 for Windows) now available
NEWS 21 JUN 13 RUSSIAPAT: New full-text patent database on STN
NEWS
     22 JUN 13
                FRFULL enhanced with patent drawing images
     23 JUN 20
NEWS
                MEDICONF to be removed from STN
NEWS 24 JUN 27
                MARPAT displays enhanced with expanded G-group definitions
                and text labels
NEWS 25 JUL 01 MEDICONF removed from STN
NEWS EXPRESS
             JUNE 13 CURRENT WINDOWS VERSION IS V8.0, CURRENT
             MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP),
             AND CURRENT DISCOVER FILE IS DATED 13 JUNE 2005
NEWS HOURS
             STN Operating Hours Plus Help Desk Availability
NEWS INTER
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             Welcome Banner and News Items
NEWS LOGIN
NEWS PHONE
             Direct Dial and Telecommunication Network Access to STN
NEWS WWW
             CAS World Wide Web Site (general information)
```

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FILE 'HOME' ENTERED AT 12:41:49 ON 06 JUL 2005

=> file reg COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION 0.21 0.21

FULL ESTIMATED COST

FILE 'REGISTRY' ENTERED AT 12:41:56 ON 06 JUL 2005 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2005 American Chemical Society (ACS)

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STRUCTURE FILE UPDATES: 5 JUL 2005 HIGHEST RN 853879-48-8 DICTIONARY FILE UPDATES: 5 JUL 2005 HIGHEST RN 853879-48-8

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TSCA INFORMATION NOW CURRENT THROUGH JANUARY 18, 2005

Please note that search-term pricing does apply when conducting SmartSELECT searches.

* The CA roles and document type information have been removed from *

* the IDE default display format and the ED field has been added, *

* effective March 20, 2005. A new display format, IDERL, is now *

* available and contains the CA role and document type information. *

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. For more information enter HELP PROP at an arrow prompt in the file or refer to the file summary sheet on the web at: http://www.cas.org/ONLINE/DBSS/registryss.html

=>

L1 STRUCTURE UPLOADED

=> s 11

SAMPLE SEARCH INITIATED 12:47:47 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED - 154 TO ITERATE

100.0% PROCESSED 154 ITERATIONS SEARCH TIME: 00.00.01

29 ANSWERS

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

PROJECTED ITERATIONS: 2336 TO 3824
PROJECTED ANSWERS: 257 TO 903

L2 29 SEA SSS SAM L1

=> d 112 1 5

'LL2' IS NOT A VALID FORMAT FOR FILE 'REGISTRY'

The following are valid formats:

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Substance information can be displayed by requesting individual
fields or predefined formats. The predefined substance formats
are: (RN = CAS Registry Number)
REG
      - RN
SAM
       - Index Name, MF, and structure - no RN
       - All substance data, except sequence data
FIDE
       - FIDE, but only 50 names
IDE
SQIDE - IDE, plus sequence data
SQIDE3 - Same as SQIDE, but 3-letter amino acid codes are used
       - Protein sequence data, includes RN
SQD3
       - Same as SQD, but 3-letter amino acid codes are used
SON
      - Protein sequence name information, includes RN
       - Table of calculated properties
CALC
EPROP - Table of experimental properties
      - EPROP and CALC
PROP
Any CA File format may be combined with any substance format to
obtain CA references citing the substance. The substance formats
must be cited first. The CA File predefined formats are:
ABS -- Abstract
APPS -- Application and Priority Information
BIB -- CA Accession Number, plus Bibliographic Data
CAN -- CA Accession Number
CBIB -- CA Accession Number, plus Bibliographic Data (compressed)
IND -- Index Data
IPC -- International Patent Classification
PATS -- PI, SO
STD -- BIB, IPC, and NCL
IABS -- ABS, indented, with text labels
IBIB -- BIB, indented, with text labels
ISTD -- STD format, indented
OBIB ----- AN, plus Bibliographic Data (original)
OIBIB ----- OBIB, indented with text labels
SBIB ----- BIB, no citations
SIBIB ----- IBIB, no citations
The ALL format gives FIDE BIB ABS IND RE, plus sequence data when
it is available.
The MAX format is the same as ALL.
The IALL format is the same as ALL with BIB ABS and IND indented,
with text labels.
For additional information, please consult the following help
messages:
HELP DFIELDS -- To see a complete list of individual display fields.
HELP FORMATS -- To see detailed descriptions of the predefined formats.
ENTER DISPLAY FORMAT (IDE):end
=> d his
     (FILE 'HOME' ENTERED AT 12:41:49 ON 06 JUL 2005)
     FILE 'REGISTRY' ENTERED AT 12:41:56 ON 06 JUL 2005
               STRUCTURE UPLOADED
L1
L2
            29 S L1
```

≈> d 12 1 5 29

L2 ANSWER 1 OF 29 REGISTRY COPYRIGHT 2005 ACS on STN

RN 688307-45-1 REGISTRY

ED Entered STN: 01 Jun 2004

CN 2H-Pyran-4-carboxamide, tetrahydro-N-hydroxy-4-[[4-[4-(4-(trifluoromethoxy)phenyl]-1-piperazinyl]sulfonyl]- (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C17 H22 F3 N3 O6 S

CI COM

SR CA

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L2 ANSWER 5 OF 29 REGISTRY COPYRIGHT 2005 ACS on STN

RN 622395-09-9 REGISTRY

ED Entered STN: 01 Dec 2003

CN 1,4-Piperidinedicarboxamide, 4-[[4-(4-butoxyphenyl)-1-piperazinyl]sulfonyl]-N4-hydroxy-N1,N1-bis(1-methylethyl)- (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C27 H45 N5 O6 S

CI COM

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1907 TO DATE)

1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L2 ANSWER 29 OF 29 REGISTRY COPYRIGHT 2005 ACS on STN

RN 210915-73-4 REGISTRY

ED Entered STN: 06 Sep 1998

CN 2-Piperidinecarboxamide, 1-[[4-(4-chloro-2-methylphenyl)-1-piperazinyl]sulfonyl]-N-hydroxy- (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C17 H25 Cl N4 O4 S

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- 2 REFERENCES IN FILE CA (1907 TO DATE)
- 2 REFERENCES IN FILE CAPLUS (1907 TO DATE)

=> d his

(FILE 'HOME' ENTERED AT 12:41:49 ON 06 JUL 2005)

FILE 'REGISTRY' ENTERED AT 12:41:56 ON 06 JUL 2005

L1 STRUCTURE UPLOADED

L2 29 S L1

=> s l1 sss full

FULL SEARCH INITIATED 12:48:39 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 3033 TO ITERATE

100.0% PROCESSED 3033 ITERATIONS

ITERATIONS 500 ANSWERS

SEARCH TIME: 00.00.01

L3 500 SEA SSS FUL L1

=> save 13

ENTER NAME OR (END):ten618288/a

ANSWER SET L3 HAS BEEN SAVED AS 'TEN618288/A'

=> file caplus

COST IN U.S. DOLLARS

SINCE FILE TOTAL

ENTRY SESSION

171.58 171.79

FULL ESTIMATED COST

FILE 'CAPLUS' ENTERED AT 12:49:13 ON 06 JUL 2005 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2005 AMERICAN CHEMICAL SOCIETY (ACS)

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FILE COVERS 1907 - 6 Jul 2005 VOL 143 ISS 2 FILE LAST UPDATED: 5 Jul 2005 (20050705/ED)

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This file contains CAS Registry Numbers for easy and accurate substance identification.

=> s 13

.4 7 L3

=> d l4 1-7 bib abs fhitstr

L4 ANSWER 1 OF 7 CAPLUS COPYRIGHT 2005 ACS on STN

Full Text

- AN 2005:409509 CAPLUS
- DN 142:463765
- TI Preparation of piperidinyl- and piperazinylsulfonylmethyl hydroxamic acids and their use as protease inhibitors
- IN Brown, David L.; Grapperhaus, Margaret L.; Kassab, Darren J.; Massa, Mark A.; Mcdonald, Joseph J.; Mullins, Patrick B.; Rico, Joseph G.; Schmidt, Michelle A.
- PA Pharmacia Corporation, USA
- SO PCT Int. Appl., 644 pp. CODEN: PIXXD2
- DT Patent
- LA English

FAN.CNT 1

	PATENT	NO.			KIN	D	DATE		i	APPL	ICAT	ION I	NO.		D	ATE	
ΡI	WO 200	0425	21		A2		2005	0512	1	WO 2	004-	US36	666		2	0041	103
	W:	ΑE,	AG,	AL,	AM,	ΑT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BW,	BY,	ΒZ,	CA,	CH,
		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,
		GΕ,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	ΚP,	KR,	ΚZ,	LC,
		LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NA,	NI,
		NO,	ΝZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,
		ТJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	ZW
	RW	BW,	GH,	GM,	KΕ,	LS,	MW,	MŻ,	NA,	SD,	SL,	SZ,	TZ,	ŪĠ,	ZM,	ZW,	AM;
		ΑZ,	BY,	KG,	ΚZ,	MD,	RU,	ТJ,	TM,	ΑT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,
		EE,	ES,	FI,	FR,	GB,	GR,	ΗU,	ΙĖ,	IS,	IT,	LU,	MC,	NL,	PL,	PT,	RO,
		SE,	SI,	SK,	TR,	BF,	ΒJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,
		NE,	SN,	TD,	TG												
PRAI GI	US 2003		A		2003	1103											

HO
$$A_{1}$$
 A_{2} A_{2} A_{3} A_{4} A_{2} A_{3} A_{4} A_{5} A

AB Title compds. I [A1-2 = H, alkyl, alkoxyalkyl, etc.; Rx = halo, CN, OH, NO2, etc.; E2 = CO, COO, OCO, amino, etc.; E3 = alkyl, alkenyl, alkynyl, etc.] are prepd. For instance, 4-[[4-(5-butylpyrazin-2-yl)piperazin-1-

yl]sulfonyl]-N-(hydroxy)tetrahydro-2H-pyran-4-carboxamide•2HCl (II) is prepd. in 8 steps from 1-(tert-butoxycarbonyl)piperazine, 2-chloropyrazine, butylmagnesium chloride, bis(2-bromoethyl)ether and O-(tetrahydro-2H-pyran-2-yl)hydroxyamine. II exhibits Ki = >10,000 nM for MMP-1, 1.52 nM for MMP-2, 0.696 nM for MMP-9, 1.82 nM for MMP-13 and 4290 nM for MMP-14. I are useful for the treatment of conditions assocd. with MMP activity and/or aggrecanase activity.

IT 622386-20-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(intermediate; prepn. of piperidinyl-and piperazinyl-sulfonylmethyl hydroxamic acids and their use as matrix metalloproteinase inhibitors)

RN 622386-20-3 CAPLUS

CN 2H-Pyran-4-carboxamide, tetrahydro-N-[(tetrahydro-2H-pyran-2-y1)oxy]-4-[[4-[4-(2,2,2-trifluoroethoxy)phenyl]-1-piperazinyl]sulfonyl]- (9CI) (CA INDEX NAME)

L4 ANSWER 2 OF 7 CAPLUS COPYRIGHT 2005 ACS on STN

Full Text

AN 2004:718284 CAPLUS

DN 141:236618

TI Inhibitors of hepatitis C virus, compositions and treatments using the same

IN Duggal, Rohit; Patick, Amy Karen; Zhao, Weidong; Herlihy, Koleen Jill; Sha, Eiann; Liu, Wei

PA Pfizer Inc., USA

SO PCT Int. Appl., 48 pp. CODEN: PIXXD2

DT Patent

LA English

FAN CNT 1

FAN.	CNT	1																
	PAT	CENT 1	NO.			KIN)	DATE		1	APPL	ICAT:	ION 1	NO.		D	ATE	
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PI	WO	2004	0735	99		A2		2004	0902	1	WO 2	004-	IB40	3		2	0040	206
	WO	2004	0735	99		A3		2004	1223									
		W:	ΑE,	ΑE,	AG,	AL,	AL,	AM,	AM,	AM,	ΑT,	ΑT,	AU,	ΑZ,	ΑZ,	BA,	BB,	BG,
			BG,	BR,	BR,	BW,	BY,	BY,	ΒZ,	ΒZ,	CA,	CH,	CN,	CN,	CO,	CO,	CR,	CR,
			CU,	CU,	CZ,	CZ,	DE,	DE,	DK,	DK,	DM,	DZ,	EC,	EC,	EE,	EE,	EG,	ES,
			ES,	FI,	FI,	GB,	GD,	GE,	GE,	GH,	GM,	HR,	HR,	HU,	HU,	ID,	IL,	IN,
			IS,	JP,	JP,	KE,	KE,	KG,	KG,	KP,	ΚP,	KP,	KR,	KR,	ΚZ,	KZ,	ΚZ,	LC,
			LK,	LR,	LS,	LS,	LT,	LU,	LV,	MA,	MD,	MD,	MG,	MK,	MN,	MW,	MX,	MX,
			MZ,	MZ,	NA,	NI												
		RW:	BW,	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	ŪĠ,	ZM,	ZW,	AT,	BE,
			BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HU,	ΙE,	IT,	LU,
			MC,	NL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,
			GQ,	GW,	ML,	MR,	ΝE,	SN,	TD,	TG,	BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,

GQ, GW, ML, MR, NE, SN, TD, TG

Р

US 2004229817 A1 20041118

20041118 US 2004-782679 20030218 20040218

PRAI US 2003-448253P

MARPAT 141:236618

AB The invention relates to methods of inhibiting HCV viral replication activity comprising contacting an HCV polymerase with a therapeutically effective amt. of a hydroxamate MMP inhibitor, and compn. comprising the same.

IT 210915-19-8

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(inhibitors of hepatitis C virus)

RN 210915-19-8 CAPLUS

CN 2-Piperidinecarboxamide, 1-[[4-(4-chlorophenyl)-1-piperazinyl]sulfonyl]-N-hydroxy- (9CI) (CA INDEX NAME)

L4 ANSWER 3 OF 7 CAPLUS COPYRIGHT 2005 ACS on STN

Apps

Full Text

AN 2003:875282 CAPLUS

DN 139:364961

- TI Preparation of piperidinyl-and piperazinyl-sulfonylmethyl hydroxamic acids and their use as protease inhibitors
- IN Barta, Thomas E.; Becker, Daniel P.; Bedell, Louis J.; Boehm, Terri L.; Brown, David L.; Carroll, Jeffery N.; Chen, Yiyuan; Fobian, Yvette; Freskos, John N.; Gasiecki, Alan F.; Grapperhaus, Margaret; Heintz, Robert M.; Hockerman, Susan L.; Kassab, Darren J.; Khanna, Ish Kumar; Kolodziej, Stephen A.; Massa, Mark; Mcdonald, Joseph; Mischke, Brent V.; Mischke, Deborah A.; Mullins, Patrick B.; Nagy, Mark; Norton, Monica B.; Rico, Joseph G.; Schmidt, Michelle A.; Stehle, Nathan W.; Talley, John J.; Vernier, William F.; Villamill, Clara I.; Wang, Lijuan Jane; Wynn, Thomas A.
- PA Pharmacia Corporation, USA; et al.
- SO PCT Int. Appl., 819 pp.

CODEN: PIXXD2

DT Patent

LA English

FAN.CNT 1

	O	-																	
	PAT	ENT I	NO.			KIN)	DATE		1	APPL	ICAT:	ION 1	NO.		D	ATE		
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PΙ	WO	2003	0912	47		A2		2003	1106	1	WO 2	003-1	US13	123		2	00304	425	
	WO	2003	0912	47		A 3		2004	0115										
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BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
     CA 2483314
                                20031106
                                             CA 2003-2483314
                                                                    20030425
                          AA
    US 2005009838
                          A1
                                20050113
                                             US 2003-618288
                                                                    20030425
    EP 1501827
                          A2
                                             EP 2003-718529
                                20050202
                                                                    20030425
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK
    BR 2003009671
                          Α
                                20050503
                                            BR 2003-9671
                                                                    20030425
PRAI US 2002-375598P
                          Ρ
                                20020425
    US 2002-380713P
                          Р
                                20020515
    US 2002-392021P
                                20020627
                          P
    WO 2003-US13123
                          W
                                20030425
os
    MARPAT 139:364961
GI
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* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Title compds. I [A1 and A2 together with the C to which they are bonded join to form (un)substituted-heterocyclyl or -carbocyclyl, or A1 and A2 are independently selected from H, alkyl, alkoxyalkyl, alkenyl, alkynyl, etc.; Rx = H, halo, CN, OH, NO2, alkyl, alkenyl, alkoxy, alkoxyalkyl, heterocyclyl, etc.; Y = N, CH, or CRx; E1 = (un)substituted heteroaryl; E2 = 0, CO, C(0)0, OC(0), bond, S, etc.; E3 = halo, CN, (un)substitutedalkyl, -alkenyl, -alkynyl, -heterocyclyl, heterocyclylalkyl, etc.] and their pharmaceutically acceptable salts are prepd. and disclosed as protease inhibitors. Thus, e.g., II·HCl was prepd. with piperazine ring formation occurring via cyclization of 2,2,2-trifluoroethoxyaniline (prepn. given) with N,N-di(2-chloroethyl)methylsulfonamide (prepn. given) to provide piperazinyl intermediate III which was converted in five addnl. steps to the desired product. This invention is directed generally to proteinase (also known as 'protease') inhibitors, and more particularly, inhibitors of matrix metalloproteinase (also known as 'matrix metalloprotease' or 'MMP') activity and/or aggrecanase activity. assays to det. inhibition consts. (Ki) against MMP-1, MMP-2, MMP-9, MMP-13 and MMP-14, I possessed values ranging from 0.13->10,000. This invention also is directed to compns. of such hydroxamic acids, intermediates for the syntheses of such hydroxamic acids, methods for making such hydroxamic acids, and methods for treating conditions (particularly pathol. conditions) assocd. with MMP activity and/or aggrecanase activity.

IT 622394-08-5P

RN

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(claimed compds.; prepn. of piperidinyl-and piperazinyl-sulfonylmethyl
hydroxamic acids and their use as matrix metalloproteinase inhibitors)
622394-08-5 CAPLUS

CN 2H-Pyran-4-carboxamide, tetrahydro-N-hydroxy-4-[[4-[4-[4-[(2,2,2-trifluoroethoxy)methyl]phenyl]-1-piperazinyl]sulfonyl]- (9CI) (CA INDEX NAME)

```
ANSWER 4 OF 7 CAPLUS COPYRIGHT 2005 ACS on STN
```

Full Text

- AN 2002:312012 CAPLUS
- 136:340996 DN
- ΤI Preparation of sulfamides as metalloprotease inhibitors
- Broka, Chris Allen; Campbell, Jeffrey Allen; Castelhano, Arlindo Lucas; TN Chen, Jian Jeffrey; Hendricks, Robert Than; Melnick, Michael Joseph; Walker, Keith Adrian Murray
- PA Syntex (U.S.A.) LLC, USA; Agouron Pharmaceuticals, Inc.
- SO U.S., 47 pp., Cont.-in-part of U.S. 6,143,744. CODEN: USXXAM
- DT Patent
- LA English.

FAN.CNT 2

		PENT						DATE											
ΡI		6376																9991	
	CA	2278	694			AA		1998	0730	(CA	199	98-	2278	694		19	980	114
	ΑU	9866	140			A1		1998	0818	7	ΑU	19	98-	6614	0		19	980	114
	ΑU	7301	27			B2		2001	0222										
	ΕP	9582	87			A1		1999	1124	1	ΕP	199	98-	9079	43		19	9980	114
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	US	6130	220			Α		2000	1010	τ	US	199	99-:	3696	77		19	990	805
	US	6143	744			Α		2000	1107	Ţ	US	199	99-3	3695	01		19	990	805
PRAI	US	1997	-367	14P		P		1997	0123										
	US	1997	-6220	09P		P		1997	1016										
	US	1998	-995	1		A3		1998	0121										
	US	1999	-369	501		A2		1999	0805										
	WO	1998	-EP1	В О		W		1998	0114										
os	MAI	RPAT	136:3	3409	96														

Sulfamides RCOCR1R2NR3SO2NR4R5 [R = OH, NHOH or N/O-alkyl or -aryl derivs.; R1, R2, R3 = H, alkyl, alkenyl, haloalkyl, cycloalkyl, cycloalkylalkyl, (hetero)aryl, acylalkyl, etc.; R1R2C may be a (hetero)carbocycle or R3 together with R1 or R2 form a heterocycloamino group; R4, R5 = H, alkyl, heteroalkyl, cycloalkyl, cycloalkylalkyl, aryl, (hetero)aralkyl or -aralkenyl; R4R5N may be a heterocycloamino group or R4 or R5 together with R3 forms an alkylene group (with provisos)], as individual isomers or mixts. of isomers, or their pharmaceuticallyacceptable salts or prodrugs were prepd. as inhibitors of metalloproteases. Thus, 2-(R)-[(1,2,3,4-tetrahydro- β -carbolino-2sulfonyl)aminolpropionic acid (claimed compd.) was prepd. by treating D-alanine Me ester hydrochloride with chlorosulfonyl isocyanate/2chloroethanol, reaction of the oxazolidone formed with 1,2,3,4-tetrahydro- β -carboline, and sapon. Metalloprotease and TNF- α inhibitory test data are tabulated.

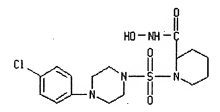
IT 210915-19-8P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of sulfamides as metalloprotease inhibitors)

RN 210915-19-8 CAPLUS

CN 2-Piperidinecarboxamide, 1-[[4-(4-chlorophenyl)-1-piperazinyl]sulfonyl]-Nhydroxy- (9CI) (CA INDEX NAME)



RE.CNT 23 THERE ARE 23 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 5 OF 7 CAPLUS COPYRIGHT 2005 ACS on STN

Full Text

AN 2000:553575 CAPLUS

DN 133:164006

TI Preparation of sulfamato hydroxamic acid metalloprotease inhibitors

IN De Crescenzo, Gary A.; Rico, Joseph G.; Boehm, Terri L.; Carroll, Jeffery
N.; Kassab, Darren J.; Mischke, Deborah A.

PA G.D. Searle and Co., USA

SO PCT Int. Appl., 628 pp.

CODEN: PIXXD2

DT Patent

LA English

FAN.CNT 1

PAN.			NO.			KIN	D	DATE		į	APP	LICAT	ION 1	NO.		D	ATE	
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			CZ,	DE,	DK,	DM,	EE,	ES,	FI,	GB,	GD	, GE,	GH,	GM,	HR,	HU,	ID,	IL,
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			SK,	SL,	ТJ,	TM,	TR,	TT,	TZ,	UA,	UG	, US,	UZ,	VN,	YU,	ZA,	ZW,	AM,
			ΑZ,	BY,	KG,	ΚZ,	MD,	RU,	TJ,	TM								
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		2362				AA						2000-						
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		2000										2000-					0000	
		6448				B1						2000-					0000: 0000:	
		2002				12 A		2002 2002				2000-		91			0000:	-
		7757		-		B2						2001-				2		
		6372						2002				2001-					0010	
		2001				A		2001				2001-					0010	
		1057				A		2002				2001-					0010	
		2001		92				2003				2001-					0010	
		6492				B1		2002				2002-					0020	
		6800				B1						2002-					0020	
		2005		80		A1		2005				2004-				2	040	708
PRAI		1999						1999										
		2000				A1		2000	0207									

WO 2000-US3061 W 20000207 US 2002-84713 A3 20020226 US 2002-262622 A3 20020930 OS MARPAT 133:164006

GΙ

AB The title compds. R20C(O)CR1R2SO2NR3aR3b (I) [wherein R1 and R2 taken together with the C to which they are attached = (un) substituted heterocyclyl or cycloalkyl; or R1 and R2 = independently H, (un) substituted (cyclo) alkyl, alkyloxylalkyl, alkylthioalkyl, alkenyl, alkynyl, aryl(alkyl), heterocyclyl(alkyl), etc.; R3a and R3b = independently H or (un) substituted alkyl, alkenyl, alkynyl, (hetero) aryl, heterocyclyl, cycloalkyl, or alkoxyalkyl; R20 = OH, alkoxyl, aryloxy, NH-OR22, or NH-OR14; R22 = selectively removable protecting group, such as 2-THP, benzyl, trisubstituted silyl, o-NO2C6H4, etc.; R14 = H, a cation, or acyl] were prepd. as selective matrix metalloproteinase (MMP) inhibitors for the treatment of various conditions, such as pathol. breakdown of connective tissue, osteoarthritis, inflammation, tumor growth, and angiogenesis. Examples include the syntheses of over 50 piperidinylsulfonyl and piperazinylsulfonyl hydroxamic acids and their intermediates. In vitro MMP assay data for I show selective inhibition of MMP-2 and MMP-13 compared to MMP-1. Some inhibition assay data for MMP-3, MMP-7, MMP-8, MMP-9, and MMP-14 are also given. Thus, II was prepd. in a multi-step sequence involving addn. of MeOC(O)Cl to 1-(methylsulfonyl)-4-(benzyloxy)piperidine (4-step prepn. given) to form the methylene sulfonamide, cycloaddn. of dibromodiethyl ether to give the THF-substituted sulfonamide, deesterification, addn. of O-(tetrahydro-2H-pyran-2-yl)hydroxylamine to form the THP hydroxamate, and deprotection to yield the desired hydroxamic acid. II inhibited MMP-1, MMP-2, and MMP-13 with IC50 values of < 10,000 nM, 7.0 nM and 20.0 nM, resp.

H

IT 287952-49-2P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(intermediate; prepn. of sulfamato hydroxamic acid metalloprotease inhibitors by cycloaddn. of dihalodialkyl ethers and amines to methylene sulfonamides followed by addn. of hydroxylamines)

RN 287952-49-2 CAPLUS

N 2H-Pyran-4-carboxamide, tetrahydro-N-[(tetrahydro-2H-pyran-2-yl)oxy]-4-[[4-[4-(trifluoromethoxy)phenyl]-1-piperazinyl]sulfonyl]- (9CI) (CA INDEX NAME)

RE.CNT 9 THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 6 OF 7 CAPLUS COPYRIGHT 2005 ACS on STN

Full Text

2000:84604 CAPLUS AN

132:141951 DN

Pharmaceutical compositions containing ACAT and MMP inhibitors for the treatment of atherosclerotic lesions

IN Bocan, Thomas Michael Andrew

Warner-Lambert Company, USA PΑ

SO PCT Int. Appl., 222 pp.

CODEN: PIXXD2

DTPatent

English LΑ

FAN.		_																
		TENT						DATE									ATE	
ΡI	WO	2000 2000	0048	92		A2						999-					9990	618
	WO					_					CN	CU,	CZ	22	CD	CE	מע	шп
		VV :																
			-	-		-		•	-	-		LR,	-		-	-		-
						•		•	•			TT,	UA,	05,	04,	VIV,	10,	ΔA,
		DW.	•	•	•	•	•	MD,	•	•		OM	λm	שמ	CIT	CV	DE	DV
		RW:		•			•	•	•	•	•	ZW,						-
			-	•						•		NL,	-	SE,	Br,	ы,	CF,	CG,
		0005	•	•					•			TD,		060				c10
		2335																
		9947															9990	
		9912															9990	
	· EP	1098															9990	
		R:				•			FR,	GB,	GR,	IT,	LΙ,	LU,	ΝL,	SE,	MC,	PT,
			•	SI,			•								_	_		
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	NO	2001	0002	91		Α											0010	118
	HR	2001	0000	55		A1		2002	0430		HR 2	001-	55			2	0010	119
PRAI	US	1998	-936	39P		₽		1998	0721			,						
	WO	1999	-US1	3948		W		1999	0618									

Acyl-CoA: cholesterol acyltransferase (ACAT) and matrix metalloproteinase (MMP) inhibitors are coadministered for the redn. of both the macrophage and smooth muscle cell component of atherosclerotic lesions, thus impairing the expansion of existing lesions and the development of new lesions and for the prevention of plaque rupture and the promotion of lesion regression in a mammal. The direct antiatherosclerotic potential of the combination of ACAT inhibitor, [[2,4,6-tris-(1-

methyl)phenyl]acetyl]-2,6-bis(1-methylethyl)phenyl sulfamic acid, and the HMG-CoA reductase inhibitor, simavastatin, in rabbits was studied. A tablet contained 2-(4'-bromobiphenyl-4-sulfonylamino)-3-Me butyric acid 25 ACAT compd. lactose 50, corn starch 20, and magnesium stearate 5 mg.

IT 210915-19-8

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(pharmaceutical compns. contg. ACAT and MMP inhibitors for treatment of atherosclerotic lesions)

RN 210915-19-8 CAPLUS

CN 2-Piperidinecarboxamide, 1-[[4-(4-chlorophenyl)-1-piperazinyl]sulfonyl]-N-hydroxy- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & & & & \\ & & & & & \\ & & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & \\ & & \\ & & \\ & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ &$$

L4 ANSWER 7 OF 7 CAPLUS COPYRIGHT 2005 ACS on STN

Full Text

AN 1998:498326 CAPLUS

DN 129:148991

TI Preparation of N-sulfamoylpiperidine-2-hydroxamic acids and analogs as metalloproteinase inhibitors

IN Broka, Chris Allen; Campbell, Jeffrey Allen; Castelhano, Arlindo Lucas; Chen, Jian Jeffrey; Hendricks, Robert Than; Melnick, Michael Joseph; Walker, Keith Adrian Murray

PA F. Hoffmann-La Roche A.-G., Switz.; Agouron Pharmaceuticals, Inc.

SO Ger. Offen., 84 pp.

CODEN: GWXXBX

DT Patent

LA German

FAN. CNT 2

FAN.	CNT	2																
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	WO	9832	748			A 1		1998	0730	,	WO 1	998-	EP18	0		1	9980	114
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			KP,	KR,	KZ,	LC,	ĹΚ,	LR,	LS,	LT,	LU,	LV,	MD,	MG,	MK,	MN,	MW,	MX,
			NO,	NZ,	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,
			UA,	ŪĠ,	UZ,	VN,	YU,	ZW,	AM,	ΑZ,	BY,	KG,	ΚZ,	MD,	RU,	ТJ,	TM	
		RW:	GH,	GM,	KE,	LS,	MW,	SD,	SZ,	ŪĠ,	ZW,	AT,	ΒE,	CH,	DE,	DK,	ES,	FI,
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	JP 3563411	B2	20040908			
	AT 223909	E	20020915	AT 1998-907943		19980114
	CN 1093125	В	20021023	CN 1998-803233		19980114
	PT 958287	T	20021231	PT 1998-907943		19980114
	ES 2183331	Т3	20030316	ES 1998-907943		19980114
	ZA 9800376	Α	19980723	ZA 1998-376		19980116
	IT 1298163 '	B1	19991220	IT 1998-MI91		19980120
	FR 2758559	A1	19980724	FR 1998-601		19980121
	GB 2321641	A1 .	19980805	GB 1998-1393		19980122
	GB 2321641	B2	20010401			
	ES 2136037	A1	19991101	ES 1998-113		19980122
	ES 2136037	B1	20001116		٤	
	NO 9903587	Α	19990922	NO 1999-3587		19990722
	NO 313635	B1	20021104			
	MX 9906822	Α	20000131	MX 1999-6822		19990722
PRAI	US 1997-36714P	P	19970123			
	US 1997-62209P	P	19971016			
	WO 1998-EP180	W	19980114			
os	MARPAT 129:148991			·		
GI						

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\end{array}$$
NHOH

II

AB R10COCR1R2NR3SO2NR20R21 [I; R1-R3 = H, (CO-interrupted) alkyl, heterocyclyl(alkyl), (hetero)aryl(alkyl), etc.; R1R2, R1R3, R2R3 = atoms to complete a ring; R10 = NR11OR12; R11,R12 = H or (ar)alkyl; R20,R21 = H, alkyl, (hetero)aryl[alk(en)yl], etc.; NR20R21heterocyclyl] were prepd. Thus, (R)-1-[4-(4-chlorobenzoyl)piperidine-1-sulfonyl]piperidine-2carboxylic acid was amidated by H2NOCMe3 and the product deprotected to give title compd. (R)-II. Data for biol. activity of I were given. IT 210915-19-8P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of N-sulfamoylpiperidine-2-hydroxamic acids and analogs as metalloproteinase inhibitors)

RN 210915-19-8 CAPLUS

CN 2-Piperidinecarboxamide, 1-[[4-(4-chlorophenyl)-1-piperazinyl]sulfonyl]-Nhydroxy- (9CI) (CA INDEX NAME)

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Full Text
AN
    2004:718284 CAPLUS
DN
    141:236618
    Inhibitors of hepatitis C virus, compositions and treatments using the
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IN
    Duggal, Rohit; Patick, Amy Karen; Zhao, Weidong; Herlihy, Koleen Jill;
    Sha, Eiann; Liu, Wei
PΑ
    Pfizer Inc., USA
SO
    PCT Int. Appl., 48 pp.
    CODEN: PIXXD2
DT
    Patent
LΑ
    English
FAN.CNT 1
    PATENT NO.
                        KIND
                               DATE
                                           APPLICATION NO.
                                                                  DATE
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                               20040902
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            ES, FI, FI, GB, GD, GE, GE, GH, GM, HR, HR, HU, HU, ID, IL, IN,
            IS, JP, JP, KE, KE, KG, KG, KP, KP, KP, KR, KR, KZ, KZ, KZ, LC,
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            MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN,
            GQ, GW, ML, MR, NE, SN, TD, TG, BF, BJ, CF, CG, CI, CM, GA, GN,
            GQ, GW, ML, MR, NE, SN, TD, TG
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                                           US 2004-782679
                                                                  20040218
PRAI US 2003-448253P
                         Р
                               20030218
OS MARPAT 141:236618
IT 210915-19-8 256646-40-9
    RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL
     (Biological study); USES (Uses)
        (inhibitors of hepatitis C virus)
RN
    210915-19-8 CAPLUS
    2-Piperidinecarboxamide, 1-[[4-(4-chlorophenyl)-1-piperazinyl]sulfonyl]-N-
    hydroxy- (9CI) (CA INDEX NAME)
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RN 256646-40-9 CAPLUS

CN 2-Piperidinecarboxamide, 1-[[4-(4-fluorophenyl)-1-piperazinyl]sulfonyl]-N-hydroxy- (9CI) (CA INDEX NAME)

L4 ANSWER 4 OF 7 CAPLUS COPYRIGHT 2005 ACS on STN

Full Text

AN 2002:312012 CAPLUS

DN 136:340996

TI Preparation of sulfamides as metalloprotease inhibitors

IN Broka, Chris Allen; Campbell, Jeffrey Allen; Castelhano, Arlindo Lucas; Chen, Jian Jeffrey; Hendricks, Robert Than; Melnick, Michael Joseph; Walker, Keith Adrian Murray

PA Syntex (U.S.A.) LLC, USA; Agouron Pharmaceuticals, Inc.

SO U.S., 47 pp., Cont.-in-part of U.S. 6,143,744. CODEN: USXXAM

DT Patent

LA English

FAN.CNT 2

FAN.	CNT	2															
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	NZ	3366	25			Α		2001	0427	NZ	1998-	3366	25		19	9801	14
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	US	6130	220			Α		2000	1010	US	1999-	3696	77		19	9908	05
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US 1998-9951 A3 19980121 US 1999-369501 A2 19990805 WO 1998-EP180 W 19980114 MARPAT 136:340996

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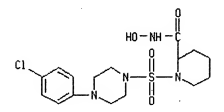
210915-73-4P 210915-75-6P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES

(prepn. of sulfamides as metalloprotease inhibitors)

RN 210915-19-8 CAPLUS

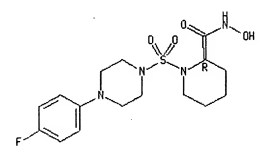
2-Piperidinecarboxamide, 1-[[4-(4-chlorophenyl)-1-piperazinyl]sulfonyl]-N-CN hydroxy- (9CI) (CA INDEX NAME)



RN210915-20-1 CAPLUS

2-Piperidinecarboxamide, 1-[[4-(4-fluorophenyl)-1-piperazinyl]sulfonyl]-N-CN hydroxy-, (2R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.





210915-32-5 CAPLUS RN

3-Isoquinolinecarboxamide, 2-[[4-(4-chlorophenyl)-1-piperazinyl]sulfonyl]-CN 1,2,3,4-tetrahydro-N-hydroxy- (9CI) (CA INDEX NAME)

RN 210915-73-4 CAPLUS

CN 2-Piperidinecarboxamide, 1-[[4-(4-chloro-2-methylphenyl)-1-piperazinyl]sulfonyl]-N-hydroxy- (9CI) (CA INDEX NAME)

RN 210915-75-6 CAPLUS

CN 2-Piperidinecarboxamide, 1-[[4-(2,3-dimethylphenyl)-1-piperazinyl]sulfonyl]-N-hydroxy- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & \\ & & \\ &$$

IT 210917-40-1P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(prepn. of sulfamides as metalloprotease inhibitors)

RN 210917-40-1 CAPLUS

CN 2-Piperidinecarboxamide, 1-[[4-(4-chlorophenyl)-1-piperazinyl]sulfonyl]-N[[(1,1-dimethylethyl)dimethylsilyl]oxy]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & \text{Me} & 0 \\ t - Bu - Si - 0 - NH - C \\ & \text{Me} & 0 \\ \hline \\ C1 & N - S - N \\ \hline \\ & N - S - N \\ \end{array}$$

RE.CNT 23 THERE ARE 23 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 5 OF 7 CAPLUS COPYRIGHT 2005 ACS on STN

Full Text

AN 2000:553575 CAPLUS

DN 133:164006

TI Preparation of sulfamato hydroxamic acid metalloprotease inhibitors

IN De Crescenzo, Gary A.; Rico, Joseph G.; Boehm, Terri L.; Carroll, Jeffery N.; Kassab, Darren J.; Mischke, Deborah A.

PA G.D. Searle and Co., USA

SO PCT Int. Appl., 628 pp.

CODEN: PIXXD2

DT Patent

LA English

FAN.CNT 1

	PA.	rent :	NO.			KIN		DATE		AI	PL	ICAT	ION 1	NO.		D.	ATE	
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			CZ,	DE,	DK,	DM,	EE,	ES,	FI,	GB, G	D,	GE,	GH,	GM,	HR,	HU,	ID,	IL,
			IN,	IS,	JP,	KE,	KG,	ΚP,	KR,	KZ, I	ď,	LK,	LR,	LS,	LT,	LU,	LV,	MA,
			MD,	MG,	MK,	MN,	MW,	MX,	NO,	NZ, E	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,
			SK,	SL,	ТJ,	TM,	TR,	TT,	TZ,	UA, U	JG,	US,	UZ,	VN,	YU,	ZA,	ZW,	AM,
	'		ΑZ,	BY,	KG,	KZ,	MD,	RU,	ТJ,	TM								
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			DK,	ES,	FI,	FR,	GB,	GR,	ΙE,	IT, I	υ,	MC,	NL,	PT,	SE,	BF,	ВJ,	CF,
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			ΙE,	SI,	LT,	LV,	FI,	RO										
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PRAI		1999				P		1999										
	US	2000	-499	276		A1		2000										
	WO	2000	-US3	061		W		2000										
	US	2002	-847	13		A3		2002										•
	US	2002	-262	622		A3		2002	0930									
os	MAI	RPAT	133:	1640						_								

IT 287952-49-2P 287953-34-8P 287953-37-1P

287953-69-9P 287954-82-9P 287954-97-6P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT

(Reactant or reagent)

(intermediate; prepn. of sulfamato hydroxamic acid metalloprotease inhibitors by cycloaddn. of dihalodialkyl ethers and amines to methylene sulfonamides followed by addn. of hydroxylamines)

RN 287952-49-2 CAPLUS

CN 2H-Pyran-4-carboxamide, tetrahydro-N-[(tetrahydro-2H-pyran-2-yl)oxy]-4-[[4-[4-(trifluoromethoxy)phenyl]-1-piperazinyl]sulfonyl]- (9CI) (CA INDEX NAME)

RN 287953-34-8 CAPLUS

CN 2H-Pyran-4-carboxamide, tetrahydro-4-[[4-(4-pentylphenyl)-1-piperazinyl]sulfonyl]-N-[(tetrahydro-2H-pyran-2-yl)oxy]- (9CI) (CA INDEX NAME)

RN 287953-37-1 CAPLUS

CN 2H-Pyran-4-carboxamide, tetrahydro-4-[(4-phenyl-1-piperazinyl)sulfonyl]-N[(tetrahydro-2H-pyran-2-yl)oxy]- (9CI) (CA INDEX NAME)

RN 287953-69-9 CAPLUS

CN 4-Piperidinecarboxamide, 4-[[4-(4-butoxyphenyl)-1-piperazinyl]sulfonyl]-1-(2-methoxyethyl)-N-[(tetrahydro-2H-pyran-2-yl)oxy]-(9CI) (CA INDEX NAME)

RN 287954-82-9 CAPLUS

CN 4-Piperidinecarboxamide, 1-(phenylmethyl)-4-[(4-phenyl-1-piperazinyl)sulfonyl]-N-[(tetrahydro-2H-pyran-2-yl)oxy]- (9CI) (CA INDEX NAME)

RN 287954-97-6 CAPLUS

CN 2H-Pyran-4-carboxamide, 4-[[4-(4-butoxy-3-methylphenyl)-1-piperazinyl]sulfonyl]tetrahydro-N-[(tetrahydro-2H-pyran-2-yl)oxy]- (9CI) (CA INDEX NAME)

IT 287952-00-5P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses)

(target compd.; prepn. of sulfamato hydroxamic acid metalloprotease inhibitors by cycloaddn. of dihalodialkyl ethers and amines to methylene sulfonamides followed by addn. of hydroxylamines)

RN 287952-00-5 CAPLUS

CN 4-Piperidinecarboxamide, N-hydroxy-1-(phenylmethyl)-4-[(4-phenyl-1-piperazinyl)sulfonyl]-, bis(trifluoroacetate) (salt) (9CI) (CA INDEX NAME)

CM 1

CRN 287951-99-9 CMF C23 H30 N4 O4 S

CM 2

CRN 76-05-1 CMF C2 H F3 O2

IT 287951-57-9P 287951-78-4P 287951-79-5P 287951-83-1P 287951-84-2P 287952-01-6P 287952-02-7P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(target compd.; prepn. of sulfamato hydroxamic acid metalloprotease

inhibitors by cycloaddn. of dihalodialkyl ethers and amines to methylene sulfonamides followed by addn. of hydroxylamines)

RN 287951-57-9 CAPLUS

CN 2H-Pyran-4-carboxamide, tetrahydro-N-hydroxy-4-[[4-[4-(trifluoromethoxy)phenyl]-1-piperazinyl]sulfonyl]-, monohydrochloride (9CI) (CA INDEX NAME)

HC1

RN 287951-78-4 CAPLUS

CN 2H-Pyran-4-carboxamide, tetrahydro-N-hydroxy-4-[[4-(4-pentylphenyl)-1-piperazinyl]sulfonyl]-, monohydrochloride (9CI) (CA INDEX NAME)

HCl

RN 287951-79-5 CAPLUS

CN 2H-Pyran-4-carboxamide, tetrahydro-N-hydroxy-4-[(4-phenyl-1-piperazinyl)sulfonyl]- (9CI) (CA INDEX NAME)

RN 287951-83-1 CAPLUS

CN 4-Piperidinecarboxamide, 4-[[4-[4-(1,1-dimethylethyl)phenyl]-1-piperazinyl]sulfonyl]-N-hydroxy-1-(2-methoxyethyl)-, monohydrochloride (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} \text{t-Bu} & & \text{CH} \text{ 2-CH} \text{ 2-OMe} \\ \hline \\ N & & \\ 0 & & \\ C & & \\ 0 & & \\ \end{array}$$

HC1

RN 287951-84-2 CAPLUS

CN 4-Piperidinecarboxamide, 4-[[4-(4-butoxyphenyl)-1-piperazinyl]sulfonyl]-N-hydroxy-1-(2-methoxyethyl)-, dihydrochloride (9CI) (CA INDEX NAME)

2 HC1

RN 287952-01-6 CAPLUS

CN 4-Piperidinecarboxamide, N-hydroxy-1-(phenylmethyl)-4-[(4-phenyl-1-piperazinyl)sulfonyl]-, dihydrochloride (9CI) (CA INDEX NAME)

2 HCl

RN 287952-02-7 CAPLUS

CN 2H-Pyran-4-carboxamide, 4-[[4-(4-butoxy-3-methylphenyl)-1-piperazinyl]sulfonyl]tetrahydro-N-hydroxy- (9CI) (CA INDEX NAME)

RE.CNT 9 THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 6 OF 7 CAPLUS COPYRIGHT 2005 ACS on STN

Full Text

AN 2000:84604 CAPLUS

132:141951 DN

TI Pharmaceutical compositions containing ACAT and MMP inhibitors for the treatment of atherosclerotic lesions

IN Bocan, Thomas Michael Andrew

PA Warner-Lambert Company, USA

SO PCT Int. Appl., 222 pp.

CODEN: PIXXD2

DTPatent

LΑ English

FAN.			NO.			KIN		DATE			APF	PLICAT	'ION	NO.		Di	ATE	
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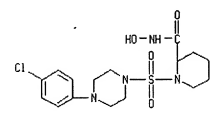
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(pharmaceutical compns. contg. ACAT and MMP inhibitors for treatment of atherosclerotic lesions)

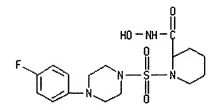
210915-19-8 CAPLUS RN

IT 210915-19-8 256646-40-9

2-Piperidinecarboxamide, 1-[[4-(4-chlorophenyl)-1-piperazinyl]sulfonyl]-N-CN hydroxy- (9CI) (CA INDEX NAME)



- RN 256646-40-9 CAPLUS
- CN 2-Piperidinecarboxamide, 1-[[4-(4-fluorophenyl)-1-piperazinyl]sulfonyl]-N-hydroxy- (9CI) (CA INDEX NAME)





L4 ANSWER 7 OF 7 CAPLUS COPYRIGHT 2005 ACS on STN

Full Text

- AN 1998:498326 CAPLUS
- DN 129:148991
- ${\tt TI}$ Preparation of N-sulfamoylpiperidine-2-hydroxamic acids and analogs as metalloproteinase inhibitors
- IN Broka, Chris Allen; Campbell, Jeffrey Allen; Castelhano, Arlindo Lucas; Chen, Jian Jeffrey; Hendricks, Robert Than; Melnick, Michael Joseph; Walker, Keith Adrian Murray
- PA F. Hoffmann-La Roche A.-G., Switz.; Agouron Pharmaceuticals, Inc.
- SO Ger. Offen., 84 pp.
 - CODEN: GWXXBX
- DT Patent
- LA German
- FAN.CNT 2

	PATENT						DATE				LICAT					ATE	
PI	DE 198															9980:	122
	CA 2278	3694			AA		1998	0730		CA	1998-	2278	694		19	9980	114
	WO 983	2748			A1		1998	0730		WO	1998-	EP18	0		19	9980:	114
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		NO,	NZ,	PL,	PT,	RO,	RU,	SD,	SE,	SG	, SI,	SK,	SL,	TJ,	TM,	TR,	TT,
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	ZA 9800 IT 1290				B1						1998-						
	FR 275										1998-						
	GB 232										1998-						
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	ES 2136037	B1	20001116		
	NO 9903587	Α	19990922	NO 1999-3587	19990722
	NO 313635	B1	20021104		
	MX 9906822	Α	20000131	MX 1999-6822	19990722
I	PRAI US 1997-36714P	P	19970123		
	US 1997-62209P	P	19971016		
	WO 1998-EP180	W	19980114		
C	OS MARPAT 129:148991				

IT 210915-19-8P 210915-20-1P 210915-32-5P

210915-73-4P 210915-75-6P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of N-sulfamoylpiperidine-2-hydroxamic acids and analogs as metalloproteinase inhibitors)

RN 210915-19-8 CAPLUS

CN 2-Piperidinecarboxamide, 1-[[4-(4-chlorophenyl)-1-piperazinyl]sulfonyl]-Nhydroxy- (9CI) (CA INDEX NAME)

RN 210915-20-1 CAPLUS

CN 2-Piperidinecarboxamide, 1-[[4-(4-fluorophenyl)-1-piperazinyl]sulfonyl]-N-hydroxy-, (2R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 210915-32-5 CAPLUS

CN 3-Isoquinolinecarboxamide, 2-[[4-(4-chlorophenyl)-1-piperazinyl]sulfonyl]-1,2,3,4-tetrahydro-N-hydroxy- (9CI) (CA INDEX NAME)

RN 210915-73-4 CAPLUS

CN 2-Piperidinecarboxamide, 1-[[4-(4-chloro-2-methylphenyl)-1-piperazinyl]sulfonyl]-N-hydroxy- (9CI) (CA INDEX NAME)

RN 210915-75-6 CAPLUS

CN 2-Piperidinecarboxamide, 1-[[4-(2,3-dimethylphenyl)-1-piperazinyl]sulfonyl]-N-hydroxy- (9CI) (CA INDEX NAME)

IT 210917-40-1P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(prepn. of N-sulfamoylpiperidine-2-hydroxamic acids and analogs as metalloproteinase inhibitors) $\,$

RN 210917-40-1 CAPLUS

CN 2-Piperidinecarboxamide, 1-[[4-(4-chlorophenyl)-1-piperazinyl]sulfonyl]-N[[(1,1-dimethylethyl)dimethylsilyl]oxy]- (9CI) (CA INDEX NAME)

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COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION FULL ESTIMATED COST 54.68 226.47

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)

SINCE FILE
ENTRY
SESSION
CA SUBSCRIBER PRICE

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FILE COVERS 1907-1966
FILE LAST UPDATED: 01 May 1997 (19970501/UP)

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